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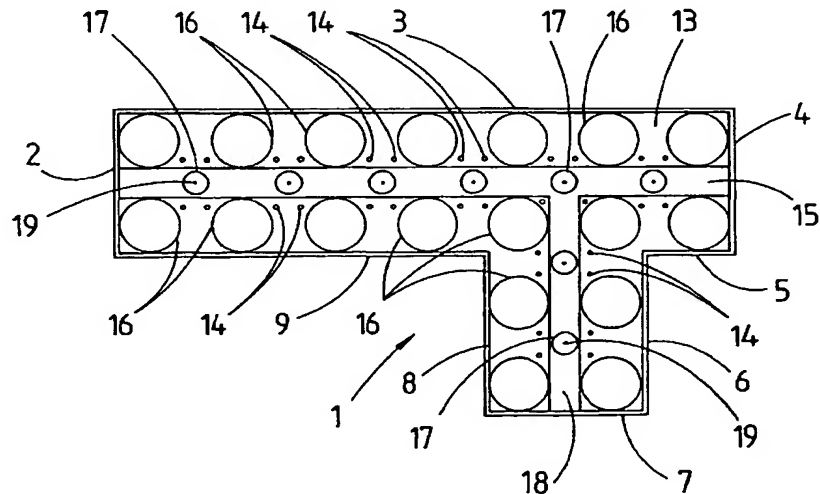
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(54) Title: METHOD OF OPERATING A FURNACE



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(57) Abstract: The method of the invention concerns operation of a furnace (1) utilising a hydrogen-rich gas as furnace fuel. The furnace (1) has a multiplicity of burners (17) for burning fuel supplied thereto. The method comprises providing ignition means for lighting a flame at at least one burner of the multiplicity of burners. An oxygen-containing gas and a combustible gas comprising a hydrocarbon gas are supplied to each of the multiplicity of burners (17) in amounts capable of forming an ignitable mixture. A flame is ignited at the predetermined one burner which is then allowed to propagate from the at least one predetermined burner to the other burners of the multiplicity of burners (17). Then the composition of the combustible gas is altered over a period of time so as to replace at least a major part of the hydrocarbon gas by a hydrogen-rich gas until a hydrogen flame is established at each of the multiplicity of burners (17).



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METHOD

The present invention is directed to a method of operating a furnace, particularly a multi-burner furnace, which utilises a hydrogen-rich gas as fuel.

When hydrogen is mixed with air in a wide range of proportions, a violent explosion can result upon ignition thereof. Moreover hydrogen has the maximum laminar burning velocity of any gas. Thus, whereas the flame speed of an acetylene flame is approximately 3.5 times that of most hydrocarbon fuels, the flame speed of a hydrogen flame is approximately 6 times higher than that of most hydrocarbons.

In certain chemical operations, particularly those involving endothermic reactions, such as steam reforming of natural gas or another hydrocarbon feedstock, it is expedient to pass the reaction mixture, for example, a mixture of the hydrocarbon feedstock and steam, through the reaction tubes of a multi-tubular reactor which are positioned in a suitable furnace and which are heated by means of a multiplicity of burners. The burners in steam reformer furnaces and other furnaces used in chemical plant operations can be supplied with any appropriate fuel, such as gas oil, natural gas, or the like. If different fuels are to be burnt, then more than one type of burner can be installed in the furnace. Often it is convenient and economical to utilise an available source of combustible waste gas as fuel for the furnace.

The reaction tubes in a steam reformer furnace typically have a nominal diameter of 5 inches (12.70 cm). They are usually mounted with their axes arranged substantially vertically and widely spaced one from another in order to allow heating by radiation and convective heating to occur. The burners can be arranged near the bottom of the furnace so that the flame extends

substantially vertically upwards, while the reactant mixture can be simultaneously fed down the catalyst-filled steam reformer tubes. In another more common arrangement the furnace is top fired. In this case the burners are mounted 5 near the top of the furnace so that the flame projects downwards into the furnace along the length of the catalyst filled, vertically mounted reformer tubes.

Other types of chemical plant which have furnaces include steam crackers for ethylene and catalytic reformers. 10 The furnaces in such forms of plant are generally top fired or side fired. Fired heaters for heating, in general, such as refinery crude heaters and vacuum unit heaters, also have multiple burners. They may burn any of a wide variety of liquid and gaseous fuels, often using more than one type of 15 burner for different fuels.

In all such furnaces the burners are normally quite widely spaced one from another and it is conventional practice to light the burners one by one with individual pilot flames or with an igniter, which is often a 20 retractable igniter, because the burners are normally spaced too far apart to allow for reliable flame propagation. To prevent burn out, the burners and pilot-light burners can be retracted into the refractory lining. Block valves are usually provided in order to allow ignition to be carried 25 out in this way and to permit maintenance of the burners. This applies also to burners which consist of a burner array, in which multiple burning points are supplied by a single supply tube or pipe off a common header. In each case it is common practice to light the burners 30 individually.

In some situations a hydrogen-rich gas is available as a waste gas stream. If, however, a hydrogen-rich gas stream is used as fuel for a furnace having a large number of

burners, there is potential for a large volume of appropriately mixed hydrogen and air to form above the burners, which will give rise to a grave risk of an explosion upon ignition of the hydrogen-rich gas stream.

5 This explosion is capable of damaging the ceramic lining to the furnace chamber or the reaction tubes or other components in the furnace and causing risk to the operators of the plant.

It is known to use a hydrogen-containing stream as a 10 fuel for a furnace. For example, it is known to utilise a methanol plant purge gas as fuel for a conventional reformer furnace. However, the purge gas stream at start up of a methanol plant is hydrogen-lean and only when the plant is 15 fully operational does a hydrogen-rich purge gas become available, by which time the burners in the furnace have already been lit. Accordingly any changeover from hydrogen-lean gas as fuel to hydrogen-rich purge gas as fuel occurs only after the burners have already been lit.

A so-called compact reformer is described in 20 International Patent Publication No. 94/29013. This has a closely packed array of reaction tubes, which are typically considerably smaller in diameter than the reaction tubes in conventional steam reformers. Thus the reaction tubes in a compact reformer typically have, for example, a nominal 25 diameter of 1½ inches (3.81 cm) in comparison with a nominal diameter of 5 inches (12.70 cm) which is typical for the reaction tubes of a conventional reformer. Moreover the reaction tubes are spaced much closer to one another in a compact reformer than in a conventional steam reformer with 30 the burners correspondingly being positioned closer to one another within the reaction tube matrix.

Since the burners are so much closer to one another in a compact reformer than in a conventional reformer furnace,

there is generally insufficient room to accommodate individual control valves for each burner fuel jet. Hence the burner fuel jets must in this case be supplied from a common manifold. Moreover, since space is limited, it is 5 hardly practical to provide multiple igniters or pilot flames and there would be an increased risk of burn out of the pilot-light fuel jets compared with conventional reformer furnaces. Autoignition would be another possibility but then it is not clear how this can be safely 10 achieved. A further possibility is to effect ignition at an outer burner of the array and then to rely on flame propagation to ignite the other burners. Although the burners in a compact reformer are close enough to permit flame propagation from one to another if conditions are 15 favourable, it is important that the correct range of velocities, fuel compositions and air:fuel ratios are used if the risks of explosions and of non-reliable ignition of all burners are to be avoided, particularly when the fuel concerned is hydrogen or a hydrogen-rich gas.

20 The present invention seeks to provide a method of igniting the burners of a furnace containing an array of closely spaced burners, such as a compact reformer, in a safe and reliable manner. In addition, it seeks to provide a method of operating a furnace with a multiplicity of 25 burners which are arranged in an array but which are not capable of individual control, in particular which are not provided with individual control valves. It further seeks to provide a method of initiating ignition in a furnace with a multiplicity of burners without using individual igniting 30 devices for each burner. The invention also seeks to provide a method permitting safe operation of a furnace having multiple burners utilising a hydrogen-rich gas as fuel, particularly during start-up of the furnace. It

further seeks to provide a method of operating a multi-burner furnace utilising a hydrogen-rich gas as fuel in which the risk of a potentially hazardous explosion is substantially obviated. An additional objective of the 5 invention is to provide a method of utilising safely the calorific value of a hydrogen-rich waste gas stream.

According to the present invention there is provided a method of operating a furnace utilising a hydrogen-rich gas as furnace fuel, the furnace having a multiplicity of 10 burners for burning fuel supplied thereto, which method comprises

- (a) providing ignition means for lighting a flame at at least one predetermined burner selected from the multiplicity of burners,
- 15 (b) supplying to each of the multiplicity of burners an oxygen-containing gas and a combustible gas comprising a hydrocarbon gas in amounts capable of forming an ignitable mixture,
- 20 (c) igniting a flame at the at least one predetermined burner,
- (d) allowing a flame to propagate from the at least one predetermined burner to the other burners of the multiplicity of burners, and
- 25 (e) altering the composition of the combustible gas over a period of time so as to replace at least a major part of the hydrocarbon gas by a hydrogen-rich gas until an at least predominantly hydrogen flame is established at each of the multiplicity of burners.

It will thus be seen that the method of the invention 30 utilises, initially, in addition to an oxygen-containing gas, a hydrocarbon gas in an amount sufficient to form an ignitable mixture. The oxygen-containing gas and the combustible gas are supplied separately to each of the

multiplicity of burners of the furnace, and then, once a flame has been lit at at least one predetermined burner selected from the multiplicity of burners, the resulting diffusion flame is allowed to propagate throughout the array 5 to each of the remaining burners. Once a suitable flame has been established at each of the multiplicity of burners of the furnace, the composition of the combustible gas is progressively adjusted so that the hydrocarbon gas is replaced by a hydrogen-rich gas, while the air and 10 combustible gas flow rates are adjusted so as to maintain a flame at each of the multiplicity of burners, thus obviating problems inherent in the direct ignition of the hydrogen-rich gas. Thereafter, once a flame has been established using the hydrogen-rich gas at each of the multiplicity of 15 burners, the flow rates of the hydrogen-rich gas can be increased to the full operational flow rates.

The method of the invention concerns operation of a furnace utilising a hydrogen-rich gas as furnace fuel. The furnace has a multiplicity of burners for burning fuel 20 supplied thereto. The method comprises providing ignition means for lighting a flame at at least one predetermined burner selected from the multiplicity of burners. An oxygen-containing gas and a combustible gas comprising a hydrocarbon gas are supplied to each of the multiplicity of 25 burners in amounts capable of forming an ignitable mixture. A flame is ignited at the at least one predetermined burner and allowed to propagate from the at least one predetermined burner to the other burners of the multiplicity of burners. Then the composition of the combustible gas is altered over 30 a period of time so as to replace at least a major part of the hydrocarbon gas by a hydrogen-rich gas until an at least predominantly hydrogen flame is established at each of the multiplicity of burners.

In the method of the invention all of the burners of the multiplicity of burners may be connected to a manifold through which the combustible gas is supplied.

During steps (a) to (e) a reduced volume flow rate of combustible gas is preferably used compared with the potential full operating flow rate for the combustible gas, if this were to be the fuel used to fire the furnace.

Similarly a reduced flow rate of hydrogen-rich gas is preferably used during steps (a) to (e) compared with that prevailing during full operation of the furnace using the hydrogen-rich gas. The flow rate of the oxygen-containing gas can also be correspondingly reduced during steps (a) to (e). Thus during step (e) the flow rate of the hydrogen-rich gas can be much lower than the full operating rate envisaged by the designer of the furnace, typically less than about 25% of that full operating flow rate and even as low as about 10% or less, for example about 5%, of the full operating flow rate. However, once a hydrogen flame or an at least predominantly hydrogen flame has been established at each of the multiplicity of burners, the rates of supply of hydrogen-rich gas and of oxygen-containing gas can be increased to the full operating flow rates. Hence the flow rate of hydrogen-rich gas during step (e) may be reduced compared with the flow rate of hydrogen-rich gas during subsequent operation of the furnace. Thus the flow rate of the hydrogen-rich gas during step (e) may be less than about 25% of the full operating flow rate of hydrogen-rich gas for which the furnace is designed.

In many cases it will suffice to ignite a flame in step (c) at a single predetermined burner of the multiplicity of burners. However, it may be more convenient or expedient to ignite a flame in step (c) at two or more predetermined burners of the multiplicity of burners.

Preferably the multiplicity of burners is arranged in an array in the furnace such that a flame ignited at the or each predetermined burner, for example a burner in an outer part of the array, can propagate from the at least one 5 predetermined burner to the other burners of the array.

In a preferred process the multiplicity of burners is mounted in a top portion of the furnace so that the flames from the multiplicity of burners extend downwards in use. Such a furnace can be, for example, a steam reformer furnace 10 having a plurality of reformer tubes, each containing a charge of a steam reforming catalyst, the reformer tubes being arranged with their axes extending in a substantially vertical direction, while the multiplicity of burners is arranged in an array in a top portion of the furnace for 15 heating the reformer tubes to a steam reforming temperature by means of flames extending downwards from the multiplicity of burners, and a reactant mixture comprising a mixture of steam and a hydrocarbon feedstock to be reformed is passed upwardly under steam reforming conditions through the heated 20 reformer tubes. In such an arrangement the method of the invention overcomes the problem that, if an attempt were to be made to ignite the hydrogen-containing gas directly, the downward flow of air might be insufficient to produce a high enough downward air velocity to overcome the natural 25 buoyancy of hydrogen, thus leading to a large envelope of hydrogen within the flammable region which may ignite in an explosive or uncontrolled way. If another gaseous hydrocarbon fuel, such as methane or natural gas, is used to 30 initiate ignition in a furnace with down firing in accordance with the method of the invention, then its higher density, narrower flammable limits, and lower burning velocity minimise the risk of explosion at the time of ignition.

Alternatively the multiplicity of burners can be mounted in a bottom portion of the furnace so that the flames from the multiplicity of burners extend upwards in operation of the furnace.

5 The ignition means for lighting a flame at the predetermined one of the multiplicity of burners can be any ignition means of known type. For example, it may comprise a piezo-electric device which produces a spark upon actuation thereof. Alternatively it may comprises an 10 electrically heated ignition element. It may comprise a pilot jet at which a pilot flame can be established prior to commencement of supply of combustible gas to the multiplicity of burners. It may be a retractable igniter device of known type.

15 Preferably the hydrocarbon gas is methane or natural gas. However, other hydrocarbon gases, such as ethane, propane, butane, or a mixture of two or more thereof, can be used, if desired, in place of or in admixture with natural gas or methane. The hydrocarbon gas can be mixed with an 20 inert gas, such as nitrogen, argon, or the like, so long as upon admixture with air or other oxygen-containing gas the resulting mixture remains combustible.

In a particularly preferred method the furnace to be operated is a steam reformer furnace used to produce 25 synthesis gas for use in an associated synthesis plant, such as a methanol synthesis plant, a Fischer Tropsch process plant, or an oxo plant for hydroformylation of an olefinic feedstock. Moreover the hydrogen-rich gas may comprise an unreacted waste gas stream from the synthesis plant.

30 The oxygen-containing gas can be oxygen, oxygen-enriched air, or air, but is preferably air. In this case the combustible gas supplied to the predetermined burner can be natural gas, while the oxygen-containing gas is air.

Typically the combustible gas and air are supplied to the multiplicity of burners in amounts sufficient to provide a mixture of about 4% by volume of natural gas and 96% by volume of air at each of the multiplicity of burners. After 5 a flame has been established at each of the multiplicity of burners, the amount of natural gas supplied to the multiplicity of burners can be gradually increased, incrementally or continuously, to provide a mixture of about 8% by volume of natural gas and 92% by volume of air at each 10 of the multiplicity of burners.

In a preferred method in step (e) the composition of the combustible gas is altered until the combustible gas substantially consists of the hydrogen-rich gas. In this case step (e) can be effected over a period of from about 1 15 second to about 15 minutes, preferably over a period of from about 2 seconds to about 5 minutes, even more preferably over a period of from about 5 seconds to about 1 minute.

In step (e) of the method of the invention the composition of the combustible gas is altered until at least 20 a major part (i.e. at least about 50%) of the hydrocarbon gas is replaced by the hydrogen-rich gas. It will normally be preferred to replace at least about 80%, and more often substantially 100%, of the hydrocarbon gas in the combustible gas of step (b) by the hydrogen-rich gas in step 25 (e).

Preferably the burners are mounted in a top portion of the furnace so that the flame from the at least one burner, or from the array of burners, extends downwards.

By operating a steam reformer furnace in accordance 30 with the method of the invention a hydrogen flame can be safely established in the furnace.

The hydrogen-rich gas can be pure hydrogen or a combustible mixture of hydrogen and one or more other gases,

such as inert gases (e.g. nitrogen, argon, and the like), or hydrocarbon gases, such as methane, ethane, propane, butane, and the like. Preferably it comprises at least about 50% by volume of hydrogen, more preferably at least about 80% by volume of hydrogen, up to about 99% by volume or more of hydrogen. When the furnace is a steam reformer furnace used to generate by steam reforming of methane or natural gas synthesis gas for the production of methanol, for the Fischer Tropsch process, or for use in an oxo process, the resulting synthesis gas contains an excess of hydrogen, as will be explained further below, in which case the hydrogen-rich gas can be the unreacted gas remaining after the subsequent synthesis step or steps.

The principal reactions that occur in a steam reformer tube are:-

- (1) $\text{CH}_4 + \text{H}_2\text{O} \rightleftharpoons \text{CO} + 3\text{H}_2$;
- (2) $\text{CO} + \text{H}_2\text{O} \rightleftharpoons \text{CO}_2 + \text{H}_2$; and
- (3) $2\text{CO} \rightleftharpoons \text{CO}_2 + \text{C}$.

As a result the resulting synthesis gas contains a $\text{H}_2:\text{CO}$ molar ratio of about 3:1 which is higher than the slightly greater than 2:1 $\text{H}_2:\text{CO}$ molar ratio required for methanol synthesis. The reactions involved in synthesising methanol from carbon monoxide and from carbon dioxide, which is generally present as a minor component of the synthesis gas mixture, are:

- (4) $\text{CO} + 2\text{H}_2 \rightleftharpoons \text{CH}_3\text{OH}$; and
- (5) $\text{CO}_2 + 3\text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} + \text{H}_2\text{O}$.

At all events, the synthesis of methanol from a synthesis gas produced by steam reforming of methane or natural gas results in a waste gas stream that is rich in hydrogen, which is suitable for use in the method of the invention. This waste gas can, if necessary, be subjected to suitable purification steps, such as pressure swing absorption, in

order to increase the hydrogen content of the gas prior to use as fuel in the furnace, the residual gas which is rich in carbon oxides being recycled to the interior of the reaction tubes of the methanol synthesis zone.

5 It will usually be preferable to preheat the combustible gas and/or the oxygen-containing gas, e.g. air, prior to supply to the furnace. Such pre-heating can be effected in conventional manner by heat exchange against a convenient source of heat, such as the flue gases from the
10 furnace. In this way the heat of combustion from the burners of the furnace is used to optimum efficiency. In such a pre-heating step the combustible gas and/or the oxygen-containing gas can be heated to a temperature in the range of from about 300°C to about 800°C.

15 In a preferred method according to the invention the preheated combustible gas is supplied, separately from the oxygen-containing gas, via a manifold to individual supply tubes feeding respective burners. These individual supply tubes are devoid of supply control valves. The oxygen-
20 containing gas is preheated by heat exchange with the reformed gases and the hot oxygen-containing gas is used to heat the outside of the individual supply tubes through which the combustible gas is supplied so as to preheat the combustible gas.

25 When it is desired to shut down a furnace burning a hydrogen-rich fuel, the hydrogen-rich gas flow stream can be switched to a flow of an equivalent amount by volume of inert gas, such as nitrogen, while maintaining the flow of oxygen-containing gas. As the inert gas replaces the
30 hydrogen-rich gas so the flame will be extinguished over a period of time. After a sufficient period of flow of the inert gas the risk of explosion in the burner assembly is removed thus avoiding a possibility of blowback. Extinction

of the flame will lead to cessation of the steam reforming reaction and cooling of the reformed gases exiting the reaction tubes. By maintaining a flow of air through the air inlet manifold, cooling of the furnace can be assisted.

5 In order that the invention may be clearly understood and readily carried into effect a preferred process in accordance with the invention will now be described, by way of example only, with reference to the accompanying drawings, wherein:-

10 Figure 1 is a top plan view of an experimental rig intended to simulate part of the burner array and tube array of a compact reformer furnace of the type disclosed in WO-A-94/29013; and

15 Figure 2 is a vertical section through the experimental rig of Figure 1.

Referring to the drawings, there is shown an experimental burner rig 1 which comprises a thermally insulated rectangular parallelipipedal box whose internal dimensions are 470 mm long x 115 mm wide by 1625 mm high, to 20 which is attached a second box 142 mm long x 115 mm wide x 1625 mm high. There is no wall between the two boxes and so the two boxes together form a box of generally offset T-section. The walls of the rig are formed by mild steel plates 2, 3, 4, 5, 6, 7, 8, and 9. The rig is open at its 25 upper end 10 but has a closed lower end formed by mild steel bottom plate 11. Above bottom plate 11 is a transverse plate 12 also made of mild steel which forms the top of an air collector box 13 and the floor of a combustion space within the rig 1. Plate 12 is pierced with 3 mm diameter 30 apertures 14 through which air for combustion can be drawn from air collector box 13 into a combustion chamber 15. All of the joints on the rig were sealed.

Within combustion chamber 15 there is mounted an array

of eighteen aluminium tubes 16 of 45 mm outside diameter equally spaced on a 70 mm square pitch. Tubes 16 are thus arranged to replicate the external shape of reformer tubes in a compact reformer furnace. Spaced between the tubes 16 and at the centre of the square pitch are placed a total of eight burner tubes 17 which have an external diameter of 19 mm and a length of 150 mm. Six burners 17 are arranged in a straight line and two are in the side branch formed by walls 6, 7 and 8. Alumina spheres (not shown) are added to a depth of 50 mm to fill the gap between tubes 16 and burner tubes 17 so that about 100 mm of each of the burner tubes 17 projects above the alumina spheres. The burner tubes 17 are each supplied from a fuel manifold box 18, which is 25 mm high, through a respective single 2 mm diameter hole 19.

15 Each burner tube is surrounded by four apertures 14 for supply of air thereto. This arrangement of burner tubes 17 and air supply apertures 14 provides, in combination with the dispersing effect of the alumina spheres, an effective distribution of air to burner tubes 17 similar to the forced air supply to an open furnace chamber containing reformer tubes.

In order to enable observation of the flames and flame propagation, a glass window 20 is installed in the wall 3 of the long side of the rig 1. The bottom of window 20 is level with the top end 21 of burner tubes 17. An 18 mm diameter hole 22 is provided in each of walls 2 and 4 and also in wall 7 with the bottom of hole 22 being level with the top end 21 of burner tubes 17. This hole 22 can be used for insertion of an oxygen-propane flame to act as pilot light to ignite the fuel from the adjacent burner tube 17 and provides additional viewing facilities. A mirror (not shown) is positioned at an angle above the open top end 10 of the rig 1 in such a way that observations can

conveniently be made, looking down into the rig 1, without the observer being subjected to heat and fumes.

The invention is further illustrated in the following Examples. In the Examples all gas flow rates are expressed 5 as l/h measured at 0°C and 760 mm Hg (101.33 kPa).

Example 1

Rig 1 was arranged to be supplied through fuel supply box 18 with substantially pure hydrogen gas as fuel and through air collector box 13 with air. The flow rates could 10 be measured using appropriate rotameters (not shown). A series of tests was carried out each lasting only a few seconds. The procedure adopted involved establishing an air flow from air supply box 13 into the combustion chamber 15 through apertures 14 and then establishing a fuel flow into 15 burner tubes 17 from fuel supply box 18 through apertures 19. An oxygen-propane pilot flame was first of all inserted into one of the ignition holes 22 so that, when hydrogen was subsequently supplied via manifold 18, a flame could be lit at the adjacent burner 17. If ignition did not occur the 20 fuel flow was stopped and the air flow was changed to a new value. The fuel was then re-supplied and ignition tried again. In some cases the adjacent burner 17 to the one lit with the oxygen-propane pilot flame would also light and propagation of flames to all burners would often occur. A 25 video recording was made of each ignition attempt. In those cases in which ignition occurred nitrogen was added to the combustion chamber 15 to extinguish the flames and the fuel supply was stopped. Without nitrogen addition blow-backs were frequently observed. The fuel supply was then re- 30 established at a new value and ignition attempted at a new flow rate as previously described. For each fuel flow rate a number of different air flow rates were tried. From the experiments it was determined that ignition and propagation

was feasible over a hydrogen flow range between 1600 and 5000 l/h with a range of excess air flow rates of from about 200% to about 400%. In other words ignition was feasible using about 10% v/v to about 15% v/v hydrogen in air
5 mixtures. However, it was observed that ignition and flame propagation using hydrogen as fuel was, in general, violent and erratic. There was a distinct "pop" as each burner was lit from the flame above each burner tube 17. It was noted that at higher fuel flow rates the release of energy was
10 greater and judged to be too violent, especially with lower excess air flow rates. At lower fuel flow rates the flame above the burner tubes 17 either did not form or was too weak to provide satisfactory propagation. This was also true at higher excess air flow rates. It was noted that the
15 calculated fuel/air mixture in several of these unsatisfactory low fuel flow rates was close to the reported lower flammability limit of hydrogen in air of 4.0%.

Example 2

The procedure of Example 1 was repeated except that
20 natural gas was used as fuel. It was determined that ignition and flame propagation was feasible over a natural gas flow rate between 1450 l/h and 2900 l/h with a range of excess air flows between about 80% and about 100%. It was observed that ignition and propagation with natural gas was
25 not violent and not erratic.

Example 3

The procedure of Example 2 was repeated with an air flow of 35000 l/h and a natural gas flow of 1700 l/h such that a stable flame was established at each burner. A
30 series of tests was then performed in which the natural gas flow was replaced with hydrogen at a flow rate of 4800 l/h over about 5 seconds to about 60 seconds. There were no violent or erratic changes in the flames and no explosions

during or after the transition between the fuels.

Example 4

Rig 1 is inverted with a wire mesh added to prevent the alumina spheres from falling out. The procedure of Example 5 1 is repeated with similar results except that the flames fire downwards.

Example 5

With rig 1 inverted the procedure of Example 2 is repeated with similar results except that the flames fire 10 downwards.

Example 6

The procedure of Example 3 is repeated with rig 1 still inverted. Similar results are obtained with the flames firing downwards.

CLAIMS:

1. A method of operating a furnace utilising a hydrogen-rich gas as furnace fuel, the furnace having a multiplicity of burners for burning fuel supplied thereto, which method comprises:

(a) providing ignition means for lighting a flame at at least one predetermined burner selected from the multiplicity of burners;

10 (b) supplying to each of the multiplicity of burners an oxygen-containing gas and a combustible gas comprising a hydrocarbon gas in amounts capable of forming an ignitable mixture;

(c) igniting a flame at the at least one predetermined burner;

15 (d) allowing a flame to propagate from the at least one predetermined burner to the other burners of the multiplicity of burners; and

(e) altering the composition of the combustible gas over a period of time so as to replace at least a major part of the 20 hydrocarbon gas by a hydrogen-rich gas until an at least predominantly hydrogen flame is established at each of the multiplicity of burners.

2. A method according to claim 1, wherein the multiplicity of burners is arranged in an array in the furnace such that 25 a flame ignited at the at least one predetermined burner can propagate to each of the other burners of the array.

3. A method according to claim 1 or claim 2, wherein the multiplicity of burners is mounted in a top portion of the furnace so that the flames from the multiplicity of burners 30 extend downwards in operation of the furnace.

4. A method according to claim 1 or claim 2, wherein the multiplicity of burners is mounted in a bottom portion of the furnace so that the flames from the multiplicity of

burners extend upwards in operation of the furnace.

5. A method according to any one of claims 1 to 3, wherein the furnace is a steam reformer furnace having a plurality of reformer tubes, each containing a charge of a steam

5 reforming catalyst, the reformer tubes being arranged with their axes extending in a substantially vertical direction, wherein the multiplicity of burners is arranged in an array in a top portion of the furnace for heating the reformer tubes to a steam reforming temperature by means of flames 10 extending downwards from the multiplicity of burners, and wherein a reactant mixture comprising a mixture of steam and a hydrocarbon feedstock to be reformed is passed upwardly under steam reforming conditions through the heated reformer tubes.

15 6. A method according to any one of claims 1 to 5, wherein the hydrocarbon gas comprises methane or natural gas.

7. A method according to any one of claims 1 to 6, wherein the furnace is a steam reformer furnace used to produce synthesis gas for a downstream synthesis plant and wherein 20 the hydrogen-rich gas comprises an unreacted waste gas stream from the downstream synthesis plant.

8. A method according to claim 7, wherein the downstream synthesis plant is selected from a methanol synthesis plant, a Fischer Tropsch plant, and an oxo plant.

25 9. A method according to any one of claims 1 to 8, wherein the combustible gas supplied to the predetermined burner in step (b) is natural gas, wherein the oxygen-containing gas is air, and wherein the combustible gas and air are supplied to the multiplicity of burners in amounts sufficient to 30 provide a mixture of about 4% by volume of natural gas and about 96% by volume of air at each of the multiplicity of burners.

10. A method according to claim 9, wherein, after a flame

has been established at each of the multiplicity of burners, the amount of natural gas supplied to the multiplicity of burners is increased, incrementally or continuously, to provide a mixture of about 8% by volume of natural gas and 5 about 92% by volume of air.

11. A method according to any one of claims 1 to 10, wherein in step (e) the composition of the combustible gas is altered until the combustible gas substantially consists of the hydrogen-rich gas.

10 12. A method according to claim 11, wherein step (e) is effected over a period of from about 5 seconds to about 1 minute.

13. A method according to any one of claims 1 to 12, wherein the flow rate of hydrogen-rich gas during step (e) 15 is reduced compared with the flow rate of hydrogen-rich gas during subsequent operation of the furnace.

14. A method according to claim 13, wherein the volume flow rate of the hydrogen-rich gas during step (e) is less than about 25% of the full operating flow rate of hydrogen-rich 20 gas for which the furnace is designed.

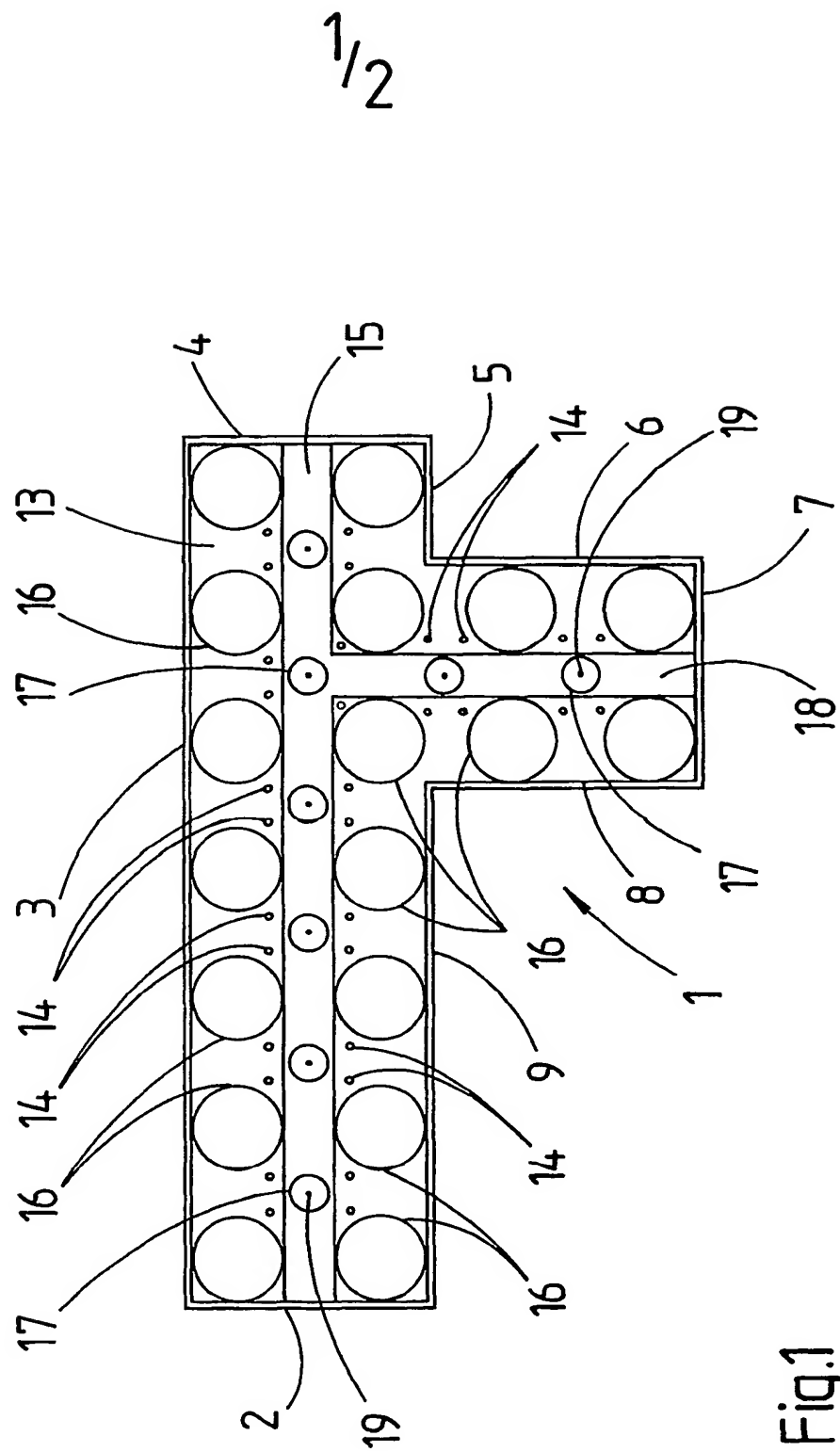


Fig.1

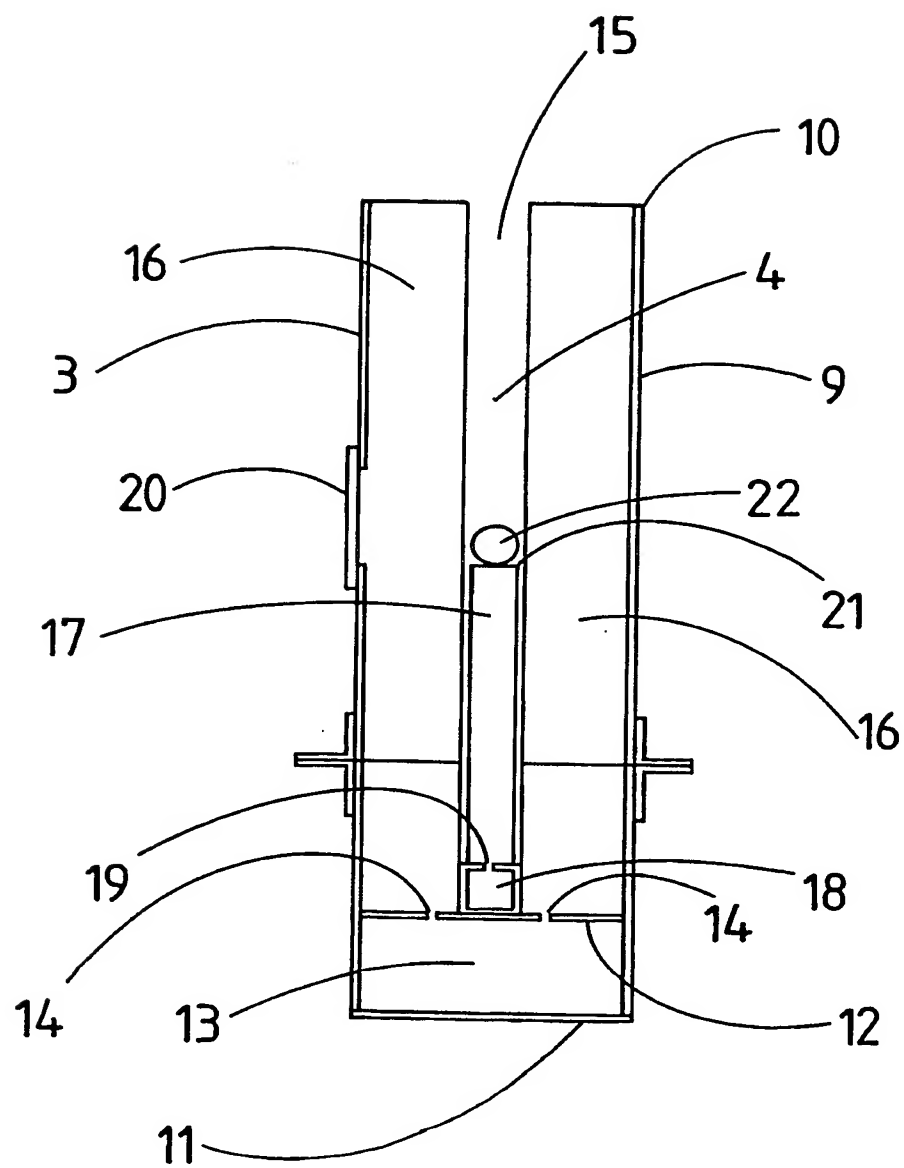
$2\frac{1}{2}$ 

Fig.2

INTERNATIONAL SEARCH REPORT

Inte Application No
PCT/GB 01/05042

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C01B3/38 F27D23/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 7 F27D C01B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)

WPI Data, PAJ, EPO-Internal

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 5 811 065 A (D.J. STERENBERG) 22 September 1998 (1998-09-22)	
A	US 6 136 279 A (H.O. STAHL) 24 October 2000 (2000-10-24)	
A	GB 1 037 094 A (THE POWER-GAS CORP) 27 July 1966 (1966-07-27)	
A	US 5 310 334 A (SPIRO SPIROS) 10 May 1994 (1994-05-10)	

Further documents are listed in the continuation of box C.



Patent family members are listed in annex.

° Special categories of cited documents :

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- "&" document member of the same patent family

Date of the actual completion of the international search

21 February 2002

Date of mailing of the international search report

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INTERNATIONAL SEARCH REPORT

Information on patent family members

Inten	Application No
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